

TRANSTEK

JUN 19 2013

Section 6 - 510(k) Summary

Date of Summary Preparation: 05/08/2013

1. Submitter's Identifications

Submitter's Name: Guangdong Transtek Medical Electronics Co., Ltd

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2. Correspondent's Identifications

Correspondent's Name: A03 Lab of BTS

Address: No.1 Fanghua Street, Hi-tech Zone, Chengdu 610041, Sichuan, China

Contact Person: Leo Wang

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3. Name of the Device

Device Classification Name: System, Measurement, Blood-Pressure, Non-invasive

Product Name: TRANSTEK Blood Pressure Monitor

Trade Name: TRANSTEK

Models: LS802-B, LS805-B, TMB-1018-BT

Classification Panel: Cardiovascular

Common/Usual Name: Automatic Blood Pressure Monitor

Product Code: DXN

Device Classification: Class II

4. The Predicate Devices

TRANSTEK, Blood Pressure Monitor, Model LS802-E, K123780

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5. Device Description

Transtek Blood Pressure Monitor, LS802-B, LS805-B and TMB-1018-BT are designed to measure the systolic and diastolic blood pressure and heartbeat rate of an individual by using a non-invasive technique in which an inflatable cuff is wrapped around the arm.

Measurement method to define systolic and diastolic pressure is similar to the auscultatory method but uses an electronic pressure sensor rather than a stethoscope and mercury manometer. The sensor converts tiny alterations in cuff pressure to electrical signals, by analyzing those signals to define the systolic and diastolic blood pressure and calculating heartbeat rate, which is a well-known technique in the market called the "Oscillometric method".

Transtek Blood Pressure Monitor is single-mounted devices of the main unit and cuff unit. ABS is used to outer housing of the main unit. The preformed cuff unit, which is applicable to arm circumference approximately between 22cm and 42cm, includes the inflatable bladder and nylon shell. The device consists of the microprocessor, the pressure sensor, the operation keys, the pump, the electromagnetic deflation control valve and the LCD. The subject device is powered by four AA/AAA alkaline batteries or by a DC 6V 400mA adapter.

The device also compares the longest and the shortest time intervals of detected pulse waves to mean time interval and displays a warning signal with the reading to indicate the detection of irregular heartbeat when the difference of the time intervals is over 25%.

Transtek Blood Pressure Monitor LS802-B, LS805-B and TMB-1018-BT embed a Bluetooth module that allows it to connect with nearby BT receiving terminal. Once measurement is over, the LCD of device displays results. And the device will start to transmit data to the paired BT terminal. Thus users can receive, display, and storage measurement data of Transtek Blood Pressure Monitor unit through their terminal devices that embedded BT module.

6. Intended Use of Device

Transtek Blood Pressure Monitor LS802-B, LS805-B and TMB-1018-BT are digital monitors intended for use in measuring blood pressure and heartbeat rate in adult patient population with arm circumference ranging from 22 cm to 42 cm (about 9 - 17 inches).

These devices detect the appearance of irregular heartbeats during measurement and give a warning signal with readings.

The Blood Pressure Monitor compares average blood pressure results to pre-established AHA (American Heart Association) hypertension guideline of 135/85 mmHg.

Transtek Blood Pressure Monitor, LS802-B, LS805-B and TMB-1018-BT are not intended to be a diagnostic device. Contact your physician if hypertensive values are indicated.

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7. Design Control Activities and Performance Tests Summary

Design control activities for this modification have been implemented and performance tests of modified devices have been done. These performance tests, risk management, and design verification tests provide demonstration that the differences do not raise any new questions of safety and effectiveness.

LS802-B, LS805-B and TMB-1018-BT conform to the following standards:

ISO 14971, Risk management to medical devices

AAMI/ANSI/IEC 80601-2-30, Safety and performance characteristics

IEC60601-1, Electrical safety; IEC60601-1-2, Electromagnetic compatibility

FCC Part 15, EMI tests of FCC Radiation & RF rules and regulations

Radio-Frequency Wireless Technology in Medical Devices, Wireless coexistence

Explanation: The wireless coexistence tests provide to demonstrate the reliability of wireless connection and wireless coexistence of BT data communication function. Meanwhile the BT data communication function does not affect blood pressure monitors' measurement function. Therefore we have not done the Clinical test.

8. Summary of Substantial Equivalence

8.1 Difference between proposed devices and the predicate device

The only significant difference between LS802-B, LS805-B and TMB-1018-BT and the predicate device is that these modified devices use Bluetooth instead of RF which the original device used.

More modification details are described in this submission.

8.2 Discussion

The Transtek Blood Pressure Monitor LS802-B, LS805-B, and TMB-1018-BT have identical indication for use, fundamental scientific technology, performance specifications, energy type, Cuff type, functions, and similar dimensional specifications, environmental specifications, software/firmware, labeling to the predicate device.

The only difference between LS802-B, LS805-B and TMB-1018-BT and the predicate device is that these modified devices' wireless data transmission function achieved by a BT module instead of original's RF module. The Bluetooth does not raise new questions of safety and effectiveness.

All required design control activities have been implemented and all applicable performance tests have been done according with demands of FDA guidance document "Non-Invasive Blood Pressure (NIBP) Monitor Guidance" FDA March 10, 1997. We found that the modified device does not create new significant risk.

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9. Conclusions

Transtek Blood Pressure Monitor LS802-B, LS805-B and TMB-1018-BT are substantially equivalent to the predicate device LS802-E, by having the identical indication for use, identical technologies, performance specification, cuff, and the similar wireless data communications, in which RF replaced by Bluetooth that does not impact the safety and effectiveness of these devices.

--- End of this section ---



June 19, 2013

Food and Drug Administration
10903 New Hampshire Avenue
Document Control Center - WO66-G609
Silver Spring, MD 20993-002

Guangdong Transtek Medical Electronics Co., Ltd
C/O Leo Wang
No.1 Fanghua Street, Hi-tech District
Chengdu, Sichuan, 610041 CH

Re: K131395

Trade/Device Name: Transtek blood pressure monitor (models: LS802-B, LS805-B, and TMB-1018-BT)

Regulation Number: 21 CFR 870.1130

Regulation Name: System, Measurement, Blood-Pressure, Non-Invasive

Regulatory Class: Class II

Product Code: DXN

Dated: May 15, 2013

Received: May 20, 2013

Dear Leo Wang:

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act) that do not require approval of a premarket approval application (PMA). You may, therefore, market the device, subject to the general controls provisions of the Act. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration. Please note: CDRH does not evaluate information related to contract liability warranties. We remind you, however, that device labeling must be truthful and not misleading.

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to additional controls. Existing major regulations affecting your device can be found in the Code of Federal Regulations, Title 21, Parts 800 to 898. In addition, FDA may publish further announcements concerning your device in the Federal Register.

Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies. You must comply with all the Act's requirements, including, but not limited to: registration and listing (21 CFR Part 807); labeling (21 CFR Part 801); medical device reporting (reporting of medical device-related adverse events) (21 CFR 803); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820); and if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.

If you desire specific advice for your device on our labeling regulation (21 CFR Part 801), please contact the Division of Small Manufacturers, International and Consumer Assistance at its toll-free number (800) 638-2041 or (301) 796-7100 or at its Internet address <http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm>. Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21 CFR Part 807.97). For questions regarding the reporting of adverse events under the MDR regulation (21 CFR Part 803), please go to <http://www.fda.gov/MedicalDevices/Safety/ReportaProblem/default.htm> for the CDRH's Office of Surveillance and Biometrics/Division of Postmarket Surveillance.

You may obtain other general information on your responsibilities under the Act from the Division of Small Manufacturers, International and Consumer Assistance at its toll-free number (800) 638 2041 or (301) 796-7100 or at its Internet address <http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm>.

Sincerely yours,

Owen P. Faris -S

for Bram D. Zuckerman, Ph.D.
Director
Division of Cardiovascular Devices
Office of Device Evaluation
Center for Devices and
Radiological Health

Enclosure

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Section 5 - Indications for Use

510(k) Number (if known):

Device Name:

Transtek Blood Pressure Monitor

Models: LS802-B, LS805-B, and TMB-1018-BT

Indications for Use:

Transtek Blood Pressure Monitor LS802-B, LS805-B, and TMB-1018-BT are digital monitors intended for use in measuring blood pressure and heartbeat rate in adult patient population with arm circumference ranging from 22 cm to 42 cm (about 9-17 inches).

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Prescription Use _____

AND/OR

Over-The-Counter Use

(Part 21 CFR 801 Subpart D)

(21 CFR 801 Subpart C)

(PLEASE DO NOT WRITE BELOW THIS LINE-CONTINUE ON ANOTHER PAGE IF NEEDED)

Concurrence of CDRH, Office of Device Evaluation (ODE)



Owen P. Faris -S
2013.06.19
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